

Chapter Ten - Informing marine spatial planning decisions with environmental DNA

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Informing marine spatial planning decisions with environmental DNA

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Abstract:

Marine management areas provide a key tool for national efforts towards sustainable development, reconciling socio-economic goals with those for biodiversity conservation. Decisions about where and when to establish spatial management areas in the oceans are currently hampered by the uncertainties of incomplete, or overly general, information about biodiversity. The analysis of environmental DNA (eDNA) provides a potentially powerful tool to overcome this lack of data in the future. Here we present directions to develop robust approaches to integrate eDNA and spatial planning processes, aiming to provide guidance to underpin tool development.

The potential of eDNA use in conservation is widely recognised, although direct applications almost exclusively focus on detection of invasive or threatened species and not spatial management decisions. The implementation of broader interaction between the fields of conservation science and eDNA analysis could create substantial benefits to biodiversity conservation and management. In particular, eDNA analysis can provide information on biodiversity over spatial-temporal scales that are currently prohibitive in spatial planning studies.

Here, we provide an overview of how eDNA is currently used in conservation practice, in addition to understanding its limitations and benefits within the context of spatial planning. With the goal to harness rapid technological developments in both molecular and conservation sciences, we provide a horizon scan of the future of eDNA analysis and its application to inform biodiversity conservation in a rapidly changing world.

Key words: eDNA, metabarcoding, tropical regions, biomonitoring, conservation, biodiversity, marine spatial planning

1. Introduction

The management of biodiversity in tropical marine ecosystems relies to a large extent on spatial management actions, such as marine protected areas (Weeks et al., 2014, Tittensor et al., 2019), no-take zones (Russ et al., 2015), and fisheries management zones (McClanahan and Hicks, 2011). Spatial planning of these management actions plays a key role in implementing objective-driven transparent prioritisations (Wilson et al., 2007, Carvalho et al., 2017), often as part of national commitments to sustainable development goals and Convention of Biological Diversity (CBD) agreements. However, spatial planning for biodiversity conservation often fails to represent biodiversity patterns and associated bio-physical processes, because up-to-date spatial databases comprising the required data are often insufficient and lack spatial, temporal, and taxonomic coverage, particularly in resource-poor tropical developing countries. The representation of biodiversity in such spatial plans often relies on very broad proxies such as ecosystem extent (Green et al., 2009, Beger et al., 2010, Makino et al., 2015), habitat type (Grantham et al., 2013, Makino et al., 2015, Boon and Beger, 2016), or bioregions (Fernandes et al., 2005) and only rarely are there attempts to capture ecosystem condition (Vercammen et al., 2019). Given the global-scale declines observed in tropical marine ecosystems (Unsworth et al., 2016, Hughes et al., 2017), up-to-date and high-resolution biodiversity data play a pivotal role in implementing rapid management responses to the climate crisis. One potential solution to these challenges lies with the emergence of environmental DNA (eDNA) technologies, defined as “genetic material obtained directly from environmental samples (soil, sediment, water, etc.) without any obvious signs of biological source material” (Thomsen and Willerslev, 2015) (Box 1).

Spatial planning evaluates the trade-offs between protecting biodiversity features and ensuring socio-economic sustainability in a transparent, quantitative, and repeatable manner (Margules and Pressey, 2000, Wilson et al., 2007, Carvalho et al., 2017, Kukkala and Moilanen, 2013). Such planning supports conservation management in efficiently reaching specific objectives by integrating multiple ecological and socio-economic variables to decide which areas should receive different conservation treatments. Importantly, effective spatial management decisions rely on carefully developed conservation objectives that are quantifiable and SMART (Specific Measurable Achievable Relevant Time-bound,

(Maxwell et al., 2015). Key principles include connectivity (e.g. considering metapopulation dynamics and larval dispersal, (Makino et al., 2015), adequacy (e.g. protecting enough of each species/ habitat, and sites that support communities in good condition, (Magris et al., 2014), representation of local biodiversity, and efficiency (e.g. minimise impact on users and other costs, (Wilson et al., 2009). Therefore, uncertainties associated with the distribution of biodiversity features and supporting processes, i.e. failure to properly implement the first three principles, can lead to management decisions that are imprecise, select unsuitable areas, or create conflict between users (Game et al., 2014).

The logistical and cost restrictions associated with underwater surveys of biodiversity, even in shallow marine habitats, the lack of taxonomic expertise, and the lack of environmental predictors operating over appropriate spatiotemporal scales for species distribution modelling (e.g. contrasting 250m² survey areas vs 100km² analysis areas in common databases such as BioOracle, (Assis et al., 2018)), all currently severely hamper spatial management decisions. Whilst there has been some progress towards integrating genetic information into conservation prioritisation (Beger, Selkoe et al. 2014, Nielsen, Beger et al. 2017), this has yet to be extended to include eDNA data. eDNA could offer solutions to many of the challenges related to scale or taxonomic expertise in collecting data needed for spatial planning (Box 1). For example, eDNA could give wider geographical coverage and span broader taxonomic ranges of biodiversity than is currently possible to record (Deiner et al. 2017; Stat et al. 2017).

Arguably one of the most exciting recent tool developments in ecology is the increasing adaptation of the analysis of eDNA to answer ecosystem-level questions, moving onwards from its origins in environmental microbiology and ancient DNA studies (Clark et al., 2018). The vast potential of eDNA analysis has led to an explosion of novel, and previously unimaginable advances in the fields of ecology (e.g. Berry et al. (2019), Deiner et al. (2017a), De Vere et al. (2017)), palaeontology (e.g. Sørnsten et al. (2010), Willerslev et al. (2014)), and conservation (e.g. Weltz et al. (2017), Cilleros et al. (2019)). eDNA has a remarkably broad spatiotemporal application spanning a diversity of environmental matrices (soil, air, aquatic systems), resolution (from haplotypes that allow for the analysis of

population level structure to entire communities) and time-frames (hours to millennia) (Ruppert et al., 2019). As such, eDNA is becoming a tool of choice for the monitoring of biodiversity, including in tropical marine environments (see for example Carvalho et al. (2019), DiBattista et al. (2019), Nichols and Marko (2019), Uthicke et al. (2018), Stat et al. (2017)) (Figure 1). However, ropical eDNA studies mostly come from a few regions such as the Red Sea (DiBattista et al., 2017, Carvalho et al., 2019) and Australia (e.g. Stat et al. 2017), leaving other regions such as South East Asia or South/Central America almost unexplored (Jerde et al., 2019). The systematic, large-scale use of eDNA and spatial planning could mean an incredible step forward in addressing this bias, where less developed, high biodiversity regions that are most threatened by human impacts are least understood and protected.

Much of the current eDNA literature has focused on tool development and addressing basic ecological questions. So far, few studies have interpreted how eDNA may be used to reveal spatiotemporal patterns and processes that could support spatial conservation decisions. In addition, inconsistencies in generating eDNA data and their analysis still limit the direct comparison and integration of different eDNA datasets that would allow for comparative analyses across larger spatial areas (McGee et al., 2019). Hence, streamlining potential future developments of eDNA tools and their application in ecology to underpin and support conservation decisions and processes presents a major opportunity in molecular and conservation sciences.

Here we assess how eDNA could be applied to spatial planning and suggest best-practice guidelines of eDNA in conservation management. Further, we propose the necessary research developments for eDNA to effectively contribute to the management and conservation of biodiversity that is applicable not only to tropical marine ecosystems, but also in terrestrial, freshwater and marine conservation efforts. We provide an overview of how eDNA is currently used in conservation practice, in addition to reviewing its limitations and benefits. We then complete a horizon scan of the future of eDNA as a tool to inform biodiversity conservation.

Current use of eDNA in natural resource management and conservation

eDNA data is often implied to be of high value for conservation initiatives (e.g. Stat et al. (2017), Ruppert et al. (2019)), but very few authors define how this would work in practice, beyond suggesting improved detection rates, or extend to the population level (Rees et al., 2014, Stat et al., 2017, Ruppert et al., 2019). eDNA has been used to detect invasive species such as bullfrogs, pythons, fish, and mussels (Ficetola et al., 2007, Hunter et al., 2015, Balasingham et al., 2017, Klymus et al., 2017, Holman et al., 2019) (Figure 1). Biomonitoring of cryptic or threatened species such as sawfishes, orang-utans, and crayfish that are challenging to monitor using conventional methods have benefitted from eDNA detection (Ikeda et al., 2016, Simpfendorfer et al., 2016, Ishige et al., 2017). Stewart et al. (2017) used eDNA to record the spatiotemporal distribution of the Yangtze finless porpoise (*Neophocaena phocaenoides*) to assess the effectiveness of protected areas in China. In addition, recent eDNA work has detected pathogens that might threaten rare populations of amphibians or crayfish (Kamoroff and Goldberg, 2017, Wu et al., 2018) (Figure 1).

eDNA studies in marine environments have included sediment, water column samples at different depths, plankton tows and stomach content. Different source materials harbour different communities and can provide complementary information (Holman et al., 2019). For example, sediments harbour more information about cryptic and benthic organisms (Pearman et al., 2018) but limited information on the fish assemblage that can be retrieved from water column samples at the same site (Koziol et al., 2019). Gut contents of predators can provide important information on their feeding behaviour or the distribution of prey (Correia et al., 2017). eDNA has also been successfully applied in monitoring the diet requirements of several species including sea lions (*Neophoca cinerea*) (Berry et al., 2017), or little penguins (*Eudyptula minor*) (Deagle et al., 2010), with studies of gut contents of parasites leading to a better understanding of the population structure of their hosts (Meekan et al., 2017).

In practice, however, few studies have used eDNA for species detection over large spatial or temporal scales in highly diverse environments such as the tropics (Cilleros et al., 2019, Stat et al., 2019). Long-term studies have tracked the effects of climate change on plankton biodiversity (Berry et al., 2019) and described shark diversity over large spatial scales (Boussarie et al., 2018). Temporal studies have

149 tracked seasonality in coastal fish communities (Sigsgaard et al., 2017b) or in lake communities (Bista
150 et al., 2017). The potential applications of eDNA analyses in threatened species management and
151 conservation science might seem endless (Figure 1), but eDNA approaches have yet to be used for
152 community analyses over large spatial scales or as part of land-based or marine spatial planning.

153 *Strategically integrating eDNA analysis and spatial planning*

154 eDNA data have the potential to revolutionise access to biodiversity information throughout the spatial
155 planning process (Figure 2). Yet to truly understand this potential, we need to assess current and
156 potential technological developments in both eDNA analysis and conservation science. An important
157 component in applying eDNA to meeting many conservation objectives, is how to treat point-based
158 eDNA data for mapping across continuous land- and sea-scapes. Even with the reduced expense that
159 eDNA could bring, fine-scale sampling across large spatial areas is likely to still have high fiscal, labour,
160 and computational costs for the foreseeable future. Therefore, innovative ways of mapping eDNA will
161 be needed to meet conservation objectives. eDNA analyses result in Operational Taxonomic Units
162 (OTUs) that serve as a proxy for species (see Glossary). The resulting sampling site \times OTU matrix will
163 still not cover the full spatial extent required for spatial planning analysis, therefore necessitating either
164 interpolation between sampling sites (Beger et al., 2014, Nielsen et al., 2017), or predictive statistical
165 modelling of OTU distributions (Figure 2). Such species distribution modelling is accomplished by
166 relating occurrence, presence/absence, or abundance data to biophysical and socio-economic predictor
167 variables (Elith and Leathwick, 2009, Guisan et al., 2013, Broennimann et al., 2012). Translation from
168 eDNA data to abundance data is currently still problematic. While some studies have found positive
169 correlations between the abundance of organisms and the quantity of eDNA molecules (Takahara et al.,
170 2012), other studies have shown that eDNA quantities are dependent upon several factors including
171 age, development stage, and environmental factors (Maruyama et al. (2014), Jo et al. (2019), Robson et
172 al., 2016).

173
174 Given the potentially small spatial resolution of eDNA data and the uncertainty associated with OTU
175 assignment, new modelling methods and small-scale biophysical parameter databases will be required.

Thereafter, OTUs could be used as a proxy for species in spatial planning, and potentially be used in three main ways when defining the spatial planning objectives (Figure 2): 1) setting conservation targets for the amount of the distribution of each OTU to be protected; 2) generating broad multivariate community types for multiple taxa and setting conservation targets for these; and 3) setting (different) targets for OTUs with different distributional patterns, such as patchy vs consistently distributed. Information derived with eDNA analysis then can help evaluate current achievements of these objectives in a gap analysis (Figure 2, Vimal et al. (2011)). With repeat sampling, changes in OTU composition and relative abundance will be able to provide information on spatiotemporal community variation (Bista et al., 2017, Berry et al., 2019) and should expand our knowledge on intra and interspecific connectivity as well as population diversity especially with the developing of organelles sequencing (Adams et al., 2019). In addition, through concerted sequencing efforts, growing barcode and genomic reference databases should facilitate species-level identification of OTUs. This may provide further insight into the prediction of the distribution of species and the use of these data in spatial planning, specifically; whether the ecological or functional traits of species can be used to further refine species distribution models.

Finally, eDNA analysis can serve as a monitoring tool, where the achievement of objectives is assessed against resampled sites over a longer term (Figure 2), and where changes in management may be adopted when required as part of an adaptive management framework (Williams and Brown, 2016). The relative ease of collecting water samples for eDNA analysis from tropical marine environments will not only allow an increase in geographical scope, but also detect a higher number of species that could be monitored simultaneously, theoretically covering the entire ecosystems' diversity. A weak point of eDNA analysis is the lack of appropriately curated and extensively populated barcode reference databases, leading to incomplete taxonomic assignment, which may be particularly problematic for mega-diverse tropical marine ecosystems. Misclassification of OTUs or ASVs is linked also to the relative short length of the amplified fragments due the degree of degradation of eDNA molecules that could reduce the possibility to discriminate between closely related organisms (Porter and Hajibabaei, 2018b).

Therefore, eDNA integration into spatial planning may require setting conservation objectives and implementing management actions for taxa (e.g. OTUs) at high taxonomic resolution (e.g. family level), whilst developing better databases (Table 2) (Porter and Hajibabaei, 2018b). Currently, spatial planning for known species applies mostly to cetaceans, reptiles and fishes, for which DNA reference databases are relatively well populated. Large global databases exist for mitochondrial (mtDNA) COI markers (Porter and Hajibabaei, 2018a), but most eDNA studies on freshwater and marine fish diversity utilise mtDNA 12S rRNA markers (Collins et al., 2019), for which reference databases are significantly smaller, as COI markers amplify phytoplankton too reducing the number of reads available for fish diversity (see **Box 1** and Glossary for details on marker regions used in eDNA work). Focussing on generating reference databases across the tree of life, in particular to adequately represent the groups with key functional roles, is pivotal for the integration of eDNA results into spatial planning (Porter and Hajibabaei, 2018b). As more eDNA data are generated, similar efforts should be applied to expand the taxonomic resolution of all the main taxonomic or functional groups of organisms.

Opportunities and challenges for eDNA analysis to inform spatial planning for conservation and management

Conservation organisations and government management agencies recognise the opportunities provided by eDNA. The unparalleled scope to detect Tree of Life assemblages of entire ecosystems with eDNA opens up multiple pathways to achieve more ambitious environmental management objectives. However, specifically within the context of spatial management, the benefits and limitations of eDNA need to be made clear, so that patterns of diversity recovered from eDNA can be interpreted appropriately. Currently, the lack of a consistent framework hampers the translation of eDNA data into spatial prioritisation plans, although eDNA data could be used to meet a wide variety of conservation objectives (**Table 1**). Major eDNA applications, from single species to whole community studies, include: 1) detection of low abundance species, 2) shortening of the time required to produce data, 3) cost reduction and 4) non-invasive or non-destructive sampling (**Table 1**).

Increasingly, it is clear that not all species can be protected or saved from extinction in the face of ubiquitous anthropogenic impacts and limited conservation funds (Bottrill, et al. 2009, Beyer et al. 2018). Instead, environmental management needs to focus on maintaining functional integrity (D'agata et al., 2016). eDNA research may provide new opportunities to define ecosystem functionality (Cordier et al., 2017), and thus to develop approaches to protect these functions in a more targeted way (Sutherland et al., 2009, Sutherland et al., 2018). Rather than measuring a small subset of species (e.g. large predators, benthic cover), eDNA sampling can detect a large proportion of micro- and macrobiota within each replicate and test how OTU interactions change between samples to ultimately discover which taxa really drive ecosystem functions (Makiola et al., 2020). For example, eDNA could identify key generalist taxa that use a wide array of environments under varying impacts, and such species could be highly resilient species of value to conservation and restoration management actions. A better understanding of how species respond to varying impacts will allow managers to detect warning signs of changing ecosystems and adapt the spatial management in response.

The potential of eDNA to target the full species assemblage, analytical methods such as ecological network analysis are likely to gain importance (Table 1) (Evans et al., 2016). Network analyses can be of use in highlighting the sensitive groups that should be targeted in management or biomonitoring (Derocles et al., 2018). Co-occurrence network analysis could reveal the way in which conservation actions (or lack thereof) affect the biodiversity of entire ecosystems (Tulloch et al., 2018). Networks could also shed light on hitherto unknown mechanisms; rare species might be influenced by other taxa that are overlooked in many of the presently used survey methods. Trophic functioning of protected versus unprotected areas could be examined in new ways as eDNA offers the potential to accurately quantify producers or invertebrates at the base of the food chain as easily as the large fishes, which currently receive the bulk of research attention (Mora et al., 2011, Cinner et al., 2016, Cinner et al., 2018, Edgar et al., 2014, Martin et al., 2017). Stomach content analysis using eDNA has also been shown to be more accurate than visual analysis and can provide much needed data on trophic interactions (Jo et al., 2016). The higher taxonomic resolution that can be achieved using eDNA

compared to conventional methods can further open up the way for far more specific spatial planning than is currently possible (Table 1).

Representing and maintaining genetic connectivity between different populations can form an important conservation objective in spatial planning (Carvalho et al., 2017). Such analyses of population genetics will increasingly benefit from eDNA data collection and advances in sequencing technology as multiple individuals (Sigsgaard et al., 2017a) and/or multiple taxa haplotypes (see Glossary) (Adams et al., 2019) can be studied at the same time. Inferring information on population genetics will become easier in the nearer future with expected cost reductions in organelle sequencing (full mitochondria or nuclear markers, see Deiner et al. (2017b)) and more comprehensive reference databases. Such advances are fundamental for monitoring and will provide information to spatial planners and managers to act rapidly and to establish best practices in ecosystem management and biodiversity conservation (Boehm et al., 2017).

Guidelines: The practicalities of spatial planning with eDNA

As with any other method, the key to obtaining accurate data relevant to conservation objectives is a well thought out experimental design and appropriate sampling method(s). While questions about the spatial coverage of eDNA are not yet fully resolved, enough is known to guide sampling design. The range at which the signature of eDNA can be detected varies from 50 m to several kilometres depending on the environment and conditions, which should be reflected in the aims and design of a study (e.g. Deiner and Altermatt (2014), Jeunen et al. (2019)). In the marine environment, eDNA can discriminate between different marine habitats, even at small spatial scales (Jeunen et al., 2019). In rivers, however, eDNA can potentially travel much further, up to 12 km, depending on river flow rates and DNA degradation time (Deiner and Altermatt, 2014, Jane et al., 2015, Pont et al., 2018). Therefore, eDNA sampling design for spatial planning depends on conservation objectives and planning area. For example, if sampling effort aims to provide information on specific habitats, it is essential to get as close as possible to the source of the eDNA which best describes that habitat. If the goal is, however, to sample a large catchment or conduct initial, general biodiversity surveys, downstream sampling could be more advisable. Moreover, the number of samples to collect (i.e. replication) needs careful

consideration under both aforementioned scenarios. Based on previous studies, more replicates with smaller volumes are preferable to a larger volume with less replication (Shaw et al., 2017, Dickie et al., 2018), and high replication ($n \gg 3$) is necessary to increase the detection rates of rarer taxa (Mächler et al., 2018, Rees et al., 2014).

While the relatively rapid degeneration of DNA makes eDNA sampling an ideal method to obtain estimates of recent diversity in a given area, this in turn brings uncertainties to designing eDNA protocols for spatial planning. Sampling might have to be repeated at multiple times throughout the year or target particular seasons (e.g. spawning events, migrations, monsoon vs dry season, etc.) depending on the goal of the sampling program (De Souza et al., 2016) and conservation objectives and spatial planning time frames. Thus, a thorough ecological understanding of the study system remains essential when designing monitoring plans (see for example Bylemans et al. (2018b)). This extends to the choice of source material. eDNA in sediments can provide precise information on different taxonomic levels, but data are less precise for obtaining information over short temporal scales. In addition, samples taken from sediments and the water column can harbour different eDNA signals (Holman et al., 2019), potentially as these capture biodiversity over longer and shorter temporal scales respectively. Conversely, sampling water could be a good solution for representing relative short time frames for a wide range of species (Collins et al., 2018).

A crucial aspect to effective tropical marine resource management is also integrating uncertainties and habitat status into spatial planning (Vercammen et al 2019), particularly for occupancy uncertainty of species or habitats. eDNA studies should therefore aim to provide such information to be relevant to conservation management. Quantifying chances of false negatives or false positives is currently not common practice in eDNA studies (but see Hunter et al. (2015)); future inclusion of these metrics would greatly benefit spatial planning. The primary risks of false positives are through contamination during sampling or extraction stages, or by wrongly assigning OTUs to a certain taxonomic species. It is essential that care is taken to avoid contamination where possible (for example, by using closed filters to capture eDNA), particularly in challenging fieldwork conditions or when engaging citizen scientists (Biggs et al., 2015, Julian et al., 2019). Increased DNA degradation rates under different conditions

could increase chances of false negatives and have unintended consequences on conservation measures, such as removal of protective measures (Chadès et al., 2008).

Finally, current eDNA methods might not be suitable for all organisms of interest (e.g. Walker et al. (2017)), or existing primer sets might not be able to detect the presence of phylogenetically distinct taxa such as Syngnathidae (Nester et al. in review). A combination of multiple primers (e.g. a “universal” 18S primer set, combined with one or more targeted primers) could therefore result in more robust data on community composition and increased detection of rare species (Deiner et al. 2017a, Berry et al., 2019) without incurring prohibitive additional costs. Technological advances such as metagenomic or whole organelle sequencing will continue to improve eDNA methodology, (Porter and Hajibabaei, 2018b). Metagenomic sequencing has the potential to remove primer and PCR biases present in metabarcoding, however, its current use in eDNA analysis remains limited due to difficulties in assigning sequences to macro-organisms (Stat et al., 2017). It is therefore advised to test the methods proposed for use with novel combinations of rare or endangered species of interest to ensure efficiency.

The future of eDNA in conservation planning science

Technological innovations can provide a step-change in conservation science and practice in the face of escalating global biodiversity declines, but such technology needs to be developed in the context of well-defined conservation problems and applications (Iacona et al., 2019). Similarly, to enhance the relevance of new directions in eDNA approaches for spatial planning, it is necessary to carefully examine and standardise important features in the methodology that relate to spatial and taxonomic comprehensiveness (Table 2), ecological relevance, and SMART (Maxwell et al., 2015) implementation of spatial planning. Spatially comprehensive and real-time biodiversity data generation will be where eDNA analysis is likely to excel, with the potential to combine eDNA with autonomous sampling and machine learning to create global monitoring networks (Bohan et al., 2017). For example, Australia and New Zealand are considering using eDNA to support biological surveillance (Cristescu and Hebert, 2018). Such systems could link with app-based spatial planning (Daigle et al., 2018) or planning web-platforms (MARXAN, www.marxan.org, SEASKETCH <https://www.seasketch.org/>) that could apply pipelines to correctly apply eDNA data. Such technology supported spatial planning

also has great potential in achieving community buy-in (Game et al., 2014). Automated eDNA recording and monitoring systems could underpin new developments in dynamic ocean planning, where eDNA data could inform where, how and what is managed on short timeframes such as a week or biweekly (Hobday et al., 2011, Lewison et al., 2015, Dunn et al., 2016). Such improvements will lead to a reduced number of field scientists and bigger studies across larger spatial and temporal scales. Thus, quantifying and reporting the impacts on ecosystem and their relative services that are largely affected by climate changes, resource overexploitation or pollution at an unprecedented resolution.

As the use of eDNA in spatial planning becomes more common, scientists from disparate fields will require simple guidelines (Figure 3). As the use of eDNA in spatial planning would benefit from remote sampling and app-based application, sharing of data and especially of metadata would become fundamental. Nowadays only the raw sequences are requested to be stored in public database, accessible to everyone, while metadata that are equally important for large regional/global comparisons are often missing or incomplete. The use of Otlet-style data and sample sharing (<https://otlet.io/>) should become a good practice step in experimental setup with clear and well documented metadata upload systems (Figure 3).

Conclusions

The natural world is currently facing multiple interacting threats on an unprecedented scale that will considerably impact how human communities connect with natural resources. Adequate resource and conservation management of tropical marine ecosystems based on real-time, large-scale biodiversity is more important than ever, yet collecting and analysing such data remains challenging. It is beyond doubt that eDNA can and will play an increasingly large role in environmental research and it will likely increase the scope of future spatial planning. This review has outlined current and future possibilities on how to do this, as well as provide information on how to integrate eDNA in planning and how to avoid the most common mistakes. The rapidly developing applications of eDNA might seem daunting to non-experts, particularly since the technique is still very recent. While research gaps and methodological uncertainties exist, the method is ready to be tested for integration in spatial planning on a large scale. Effective marine spatial planning decisions depend on accurate and timely knowledge

364 of the system to be managed. eDNA provides a step-change in how we think about the availability of
365 biodiversity data, and has the potential to completely redefine the spatio-temporal context of how
366 ecological systems are managed.

367

Glossary

Molecular terminology

Environmental DNA (eDNA): DNA directly extracted from environmental samples (soil, sediment, water, etc.) without any knowledge of the original organism.

PCR (Polymerase Chain Reaction): a molecular technique that allows the exponential amplification of a target fragment/region of DNA from a mixture of DNA fragments. The desired fragment to amplify is recognized from the other fragments in the mixture by specific primers (small single strand oligonucleotides) complementary to the desired sequence. The process is based on sequential cycles of heating and cooling at specific temperature. In the first step, the double strand DNA molecules are separated into single strands by high temperatures. In the second step, temperature is lowered, and primers bind to the complementary sequences of the targeted regions of DNA. In the third step, temperature is increased to the working optimum for the polymerase enzyme. The enzyme adds nucleotides to assemble the complementary sequence of the target DNA. During a PCR reaction, the three steps are repeated several times (between 25-30 cycles) and for each cycle the quantity of amplified DNA increases exponentially.

Quantitative PCR (qPCR): Quantitative PCR is a variant of PCR. The main difference between the two is that qPCR is able to quantify how many fragments of DNA are amplified during each step in the reaction, leading to quantitative data.

High Throughput Sequencing (HTS): a technique able to determine the nucleotide composition of millions of nucleic acid sequences. Different types of sequencing are now available and include Illumina, PacBio or NanoPore. Every sequencing method uses different strategies to generate the nucleic acid sequence, for example Illumina uses fluorescent nucleotides while NanoPore uses current change when DNA strand passes through a membrane protein. For more information please see van Dijk et al. (2018).

Metabarcoding: Taxonomic identification of millions of sequences in one experiment generated by PCR amplification on eDNA samples. This is possible using one of the HTS techniques.

Metagenomics: Different from metabarcoding, metagenomics analyses do not require PCR amplification prior to sequencing. During the process, all DNA molecules are amplified together, which limits the error connected with PCR amplification.

Operational Taxonomic Unit (OTU): Sequences (reads) obtained from HTS are grouped together, to minimise the influence of PCR and sequencing error, based on threshold dissimilarity (usually 3%). OTUs clusters are generated in programs such as VSEARCH or USEARCH. They are NOT species but an approximation to species. Clustering together multiple reads will inevitably reduce the information on nucleotide variations within that OTU.

Amplicon Sequence Variance (ASV): ASV does not include grouping reads based on dissimilarity but retains all the reads that are generated by HTS after a denoising step (removing of sequencing errors and chimera). Single nucleotide variation sequences are maintained in the dataset allowing the discrimination between different haplotypes within the same species. Programs that generate ASVs are DADA2 or Deblur. For a more detailed description of the difference between OTU and ASV see Callahan et al. (2017).

Haplotypes: a group of alleles that are inherited together from a single parent, for example mitochondrial haplotypes. Haplogroup are haplotypes that shared a common ancestor with a single nucleotide polymorphism mutation.

Cytochrome oxidase I (COI): Alternatively known as COX1 or CO1, it is a mitochondrial gene that encodes the main subunit of the cytochrome c complex. It is widely used to barcode eukaryotes. The reference database for this gene is known as BOLD database.

Barcode of Life Data (BOLD): Public database of COI gene sequenced across all the tree of Life.

12S rRNA: mitochondrial gene that is used for taxonomic assignment especially for fish.

Conservation terminology

Convention on Biological Diversity (CBD): International convention signed by 168 countries which aims to conserve biological diversity, promote sustainable use of biological diversity and ensure the fair and equitable sharing of the benefits arising out of the utilization of genetic resources.

420 **Data spatialisation:** Transformation of site-specific environmental, biodiversity, and socio-economic
421 data into spatially-explicit map-based representations, typically achieved through distribution
422 modelling or interpolation.

423 **Gap Analysis:** Method used to identify problems or gaps that are likely to decrease the efficiency of
424 protected area managements. Gaps can range from exclusion of species or habitats, to missing
425 ecological processes, or problems in the management process itself.

426 **Marine spatial planning:** Framework to decide where to implement different management and
427 conservation actions by evaluating the trade-offs between protecting biodiversity features and ensuring
428 socio-economic sustainability in a transparent, quantitative, and repeatable manner.

429 **Objectives:** Quantitative specification of management goals for a certain ecosystem, habitat type, or
430 species. Objectives can be ecological, social, or economic, but should be detailed and quantifiable.

431 **SMART (spatial planning):** A conservation approach where objectives are Specific (clearly defined),
432 Measurable (specific on what will be measured and how), Achievable (realistic in light of existing
433 ecological and social conditions), Relevant (complementary to project goals), and Time-bound (clear
434 timeline).

435

Box 1. A basic guide to eDNA studies.

The analysis of microbial life using eDNA approaches has been commonplace for over 20 years, but their use to detect macro-organisms to investigate large scale ecological processes is more recent (Ficetola et al., 2007). Recently, eDNA methods have gained increasing attention as a possible alternative to survey rare or cryptic species, or to replace lethal or invasive survey techniques (Barnes and Turner, 2016, Jeunen et al., 2019). Environmental samples used in eDNA studies collect a mixture of DNA fragments originating from the various organisms present in that environment, regardless of whether these organisms are visible or morphologically identifiable in the source material. This pool of DNA is then extracted with commercial kits or other well-established protocols. It is then often necessary to amplify the amount of DNA present via Polymerase Chain Reaction technology (PCR, see Glossary), before identifying which taxa it originates from (although methods are increasingly moving away from amplification-based approaches). Whilst widely used for microorganisms, metagenomics (see Glossary) approaches are not routinely used in eDNA studies, with, to our knowledge, only one study that has attempted this method (Stat et al. (2017)). The main limitation for metagenomics is the very limited percentage of macro-organism DNA sequences that can be amplified (Stat et al. (2017)).

eDNA studies can be broadly classified into studies focussing on community composition or those that target specific organisms or even for population-level studies (Porter and Hajibabaei, 2018a). When targeting individual species, species-specific primers (employed during PCR) ensure that only the target species is amplified, with researchers utilising quantitative (or more recently digital) PCR (see Glossary) to provide estimates of biomass or cellular abundance (Porter and Hajibabaei, 2018a). This approach is common in the detection of invasive species, such as American bullfrogs (*Rana catesbeiana*) in France (Ficetola et al., 2007), the Asian Carp (*Hypophthalmichthys* sp.) in the USA (Bohmann et al., 2014), and Asian date mussels (*Arcuatula senhousia*) (Holman et al., 2019), alongside detecting threatened species; for example, the great crested newt (*Triturus cristatus*) in the UK, Endangered skates (*Zoaraja maugeana*) in Australia, or nearly extinct freshwater fish (*Misgurnus fossilis*) of conservation concern in Denmark (Biggs et al., 2015, Sigsgaard et al., 2015, Weltz et al., 2017). Currently, eDNA tools are being tested and specialised companies are already offering related

services (see for example <https://www.naturemetrics.co.uk/wildlife-services/gcn-edna/>) particularly in single-species management of invasive or threatened species.

For studies aiming to record all species present in a sample, High Throughput Sequencing (see Glossary HTS) in the form of metabarcoding (see Glossary) is currently the most commonly applied approach. Metabarcoding is the taxonomic identification of multiple species extracted from eDNA samples (Deiner et al., 2017a). Metabarcoding primers employed during PCR aim to capture broad taxonomic groups, for example, amplifying all the eukaryotes present, or targeting specific groups such as fish or crustaceans. These primers anneal to complementary sequence in the mixed pool of DNA fragments and only amplify copies of a selected genomic region that contains enough sequence information to facilitating species identification. These amplicons are then sequenced using one of the available HTS technologies (Illumina for example uses nucleotide labelled with different fluorochromes that are read by a laser while Nanopore using differences in membrane potential see van Dijk et al. (2018) for a review on HTS). The DNA sequence reads, generated via HTS, are then analysed to determine the species composition of the original sample. A series of bioinformatics tools are employed to generate ecologically-relevant data for biomonitoring and/or spatial planning. Typically, sequences are clustered in groups based on a predefined similarity threshold (OTUs; see Glossary) or left ungrouped to capture total genetic variation (ASVs; see Glossary). OTUs and/or ASVs are then matched to sequences of known taxonomic identity held in large databases, which then completes the identification of the species present.

The analysis of eDNA provides data that leverages the ability to monitor species composition and distribution in a quicker and often easier way than more traditional approaches (Bista et al., 2017, Bohan et al., 2017, Cristescu and Hebert, 2018). However, the probability of species detection via eDNA approaches, differs in many and often unknown ways; field and laboratory methods (McGee et al., 2019), sampling depths (Eilers et al., 2012, DiBattista et al., 2019) environmental substrate (Holman et al., 2019), and the chemical, physical, oceanographic and biological factors that influence eDNA degradation (Rees et al., 2014), all introduce potential biases. For example, water samples are more homogenous than sediment samples that can contain significant small-scale heterogeneities (Koziol et

al., 2019). The distance eDNA travels from a source is highly variable, ranging from 50 meters (Jerde et al., 2016) to >200 meters (Jane et al., 2015, Pont et al., 2018). Heterogeneous sources of microbial eDNA consist of different particles (mucus, skin, faeces etc.) that are transported, settled on the benthos, and resuspended in a complex and stochastic manner (Jerde et al., 2016, Shogren et al., 2017) and that may degrade differently according to their size Jo et al. (2017), Wei et al. (2018) but see the contrary in Bista et al. (2018), Bylemans et al. (2018a). Field sampling in tropical marine environments relies on many different strategies, including surface water (Cilleros et al., 2019), benthic water (Boussarie et al., 2018), and sediment samples (DiBattista et al., 2019), mirroring non-standardised sampling in the eDNA and metabarcoding fields (McGee et al., 2019). In addition, variation in laboratory procedures may influence the comparability of results across studies (McGee et al., 2019, Berry et al., 2019, Kelly et al., 2019) however the biggest variability is how the data are produced and analysed.

Illumina technology is currently most commonly used but other options are available and comparisons between different sequencing techniques can be challenging (Porter and Hajibabaei, 2018 b). One of the most debated steps in the bioinformatics workflow is sequence clustering. Researchers can cluster their sequences in OTU (see Glossary) based on a similarity threshold or treat them as ASV (see Glossary) without clustering (Deiner et al. (2017a)), Incorrect clustering can have strong effects on alpha diversity indices introducing overestimation or under estimation within the community (Pawlowski et al., 2018). Taxonomy can be assigned using a variety of programs that are based on different approaches including BLAST, MG-RAST or RDP which can lead to different outputs (Deiner et al. (2017a)). The use of different databases can similarly lead to different annotations and potential errors. Contrary to BOLD, NCBI database is not curated but recently has been demonstrated to be reliable for eDNA analysis (Leray et al., 2019) especially because error in taxonomic assignments are easier to correct as more information on biogeography are available for macro-organisms (Deiner et al. (2017a)).

We suggest the following reviews/articles for a better understanding of the above discussed possibilities and challenges that eDNA studies face. For general application of eDNA, we recommend Rees et al.

(2014), Cristescu and Hebert (2018), for better understanding of PCR and bioinformatics application on eDNA we strongly suggest Deiner et al. (2017a) and Kelly et al. (2019).

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Table 1. Potential spatial planning objectives used in spatial planning and how the use of eDNA could influence future practice for tropical marine systems, references provided where available.

Conservation objective	Conservation action	Data needed	Current challenges	eDNA opportunities	Examples of eDNA use
<i>Identify and eradicate invasive species</i>	Robinson et al. (2019)	Presence/absence of invasive species	Low detection probability	Increased detection probability + decreased cost	Holman et al. (2019)
<i>Manage rare/threatened species</i>	Stewart et al. (2017)	Presence/absence of species	Low detection probability	Increased detection probability + decreased cost	Stewart et al. (2017)
<i>Represent cryptic species</i>	DiBattista et al. (2017)	Cryptic species detection	Low detection probability	Increased detection probability + decreased cost	Holman et al. (2019)
<i>Baseline biodiversity assessment</i>	Stat et al. (2017)	Species presence data from entire assemblage	Expensive to collect fully representative samples	Can be collected in single sample	Stat et al 2017 Cilleros et al 2019
<i>Protect ecosystem functions of poorly studied taxa</i>	Costello et al 2015	Little known taxa	High taxonomic expertise required	Reference database return species / OTUs for most taxa	Not available yet
<i>Manage ecological networks</i>	Tulloch et al. 2018	Species abundance data from entire assemblage	Intensive data collection + unlikely to cover entire assemblage	Full assemblage collected + reduced sampling effort	Not available yet
<i>Manage trophic functioning</i>	Mills et al. (2011)	Invertebrate data	Low detection probability + intensive data collection	Increased detection probability + Tree of Life assemblage collection possible	Not available yet

<i>Protect ecosystem functions of microorganism (e.g. macronutrient cycling)</i>	Louca et al. 2016	Microbial functional gene	Target studies on specific gene	Single sample can provide information on microbial functionality as well as baseline biodiversity data	Delmont et al. (2018)
<i>Find priority areas for connected protected area networks</i>	Beger et al. (2014), Nielsen et al. (2017)	Population genetics of target species	Invasive + intensive sampling effort	Non-invasive + reduced sampling effort	Sigsgaard et al. (2017a)
<i>Avoid protecting exclusion areas</i>	Daigle et al. (2018)	In-depth knowledge of ecosystem function + presence/absence data target species	Not possible or not cost-effective	Increased detection probability + decreased cost	Not available yet
<i>Monitoring: Assess trends in species populations</i>	McDonald-Madden et al. (2010)	Abundance data	Low detection probability + labour intensive to get accurate abundance of target species	Not deployed yet, easier to standardise through time	Bista et al. (2017), Sigsgaard et al. (2017b)
<i>Monitoring: Assess sex and/ or age of populations</i>	Stewart et al. 2018	In-depth knowledge of the species	Invasive + intensive sampling effort	Not deployed yet	Not available yet
<i>Monitoring: Maintain large scale (global) sampling of population trends</i>	Thompson et al. (2016)	Biodiversity data large geographic scale	Extensive, standardised sampling effort + expensive	Decreased sampling cost + easier to standardise	Gilbert et al. (2014)

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1021 **Table 2. Horizon scanning table of research needs to optimise eDNA integration in spatial**
1022 **planning, references provided where available.**

Factor	Challenge	Future	(Review) References
Abiotic and biotic degradation	Very limited information on tropical environments (e.g. Temperature, pH, UV intensity)	Quantify decay rate-biophysical conditions relationships for different settings	Lopes et al. (2017), Sales et al. (2019), Valdez-Moreno et al. (2019)
Sampling and analysis method efficiency	Differences in sampling method (substrate, volume, extraction, primers) influence results.	Quantify differences associated with different sampling methods	McGee et al. (2019), Holman et al. (2019)
	Sample size and replication varies across studies	Develop hierarchical standardised protocols for multi-purpose eDNA analyses.	Dickie et al. (2018), Mächler et al. (2018), Rees et al. (2014)
	Use of negative controls (i.e. filtration (field) and extraction blank) only.	Include positive controls (i.e. a mixture of known DNA from different species and at different concentrations)	Zinger et al. (2019), Evans et al. (2017)
Bioinformatics and statistical analysis	Data analysis methods are variable, i.e. different pipelines and algorithms	Develop a robust bioinformatic pipeline that could be used across different eDNA experimental set ups	Eckert et al. (2018), Deiner et al. (2017a)
Reference databases are lacking	Lack of curated databases for taxonomic assignment More reference data required on different markers than COI Databases lack information such as geographic position or environmental variables Rate of false positive and false negative observations not captured	Better population of databases entries and improved curation, with barcodes for identified taxa, in parallel with better bioinformatics algorithms Development of occupancy models to quantify error rates (ongoing, but more depth needed) Occupancy models should take in account the hierarchical nature of the experiment and include false positives	Porter and Hajibabaei (2018a) Ficetola et al. (2015), Ficetola et al. (2016) Davis et al. (2018), Doi et al. (2019), Strickland and Roberts (2019)
Method boundaries	Turning read abundance into estimated biomass	Develop, test, and document potential biomass/ abundance eDNA methods	Contradictory results based on few species studied (Sassoubre et al., 2016, Robson et al., 2016)

Data access and compatibility	Data storage not centralised, metadata not available	Standardised metadata recording should be established and metadata appropriately deposited. Centralise eDNA data in open access databases (with solid funding) to increase use by wider community	
Unifying spatial planning with eDNA data framework and guidelines	Very little overlap in expertise between molecular ecologists and conservation scientists	Better and bigger databases that could be used for spatial planning	Beger et al. (2014), Nielsen et al. (2017), von der Heyden et al. (2014)

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1025 **Figure 1.** Application of eDNA in conservation science. eDNA has been utilised in species detection
1026 (including cryptic, threatened, rare and invasive species), in diet analysis, or to survey biodiversity in
1027 complex environments. Each of these is applicable within different management contexts (Table 1).

1028 **Figure 2.** Flowchart of the spatial planning process (modified from Pressey and Bottrill (2009)), with
1029 potential uses of eDNA at relevant stages. **Planning:** the benefits, limitations, feasibility, and cost of
1030 eDNA approaches should be carefully considered at the planning stage; **Data collection:** eDNA could
1031 be used to provide biodiversity baseline data; **Conservation objectives:** quantifiable diversity objectives
1032 could be set by using metrics generated from eDNA data; **Current achievement of objectives:** data
1033 collected with eDNA can be compared to objectives; **Evaluation:** eDNA data can be used to monitor
1034 progress on objectives.

1035 **Figure 3.** Schematic guidelines for future eDNA projects that provide biodiversity data for spatial
1036 planning.

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